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# Application for United States Letters Patent

To all whom it may concern:

Be it known that, we,

Jayalekshmy Ayyer, Pradipkumar Maheshchandra Shah and Virendra Bhikhabhai Patel

have invented certain new and useful improvements in

A NOVEL CATALYST USEFUL FOR REMOVAL OF HYDROGEN SULPHIDE FROM GAS STREAM AND ITS CONVERSION TO SULPHUR, A PROCESS FOR PREPARING SUCH CATALYST AND A METHOD FOR REMOVING OF HYDROGEN SULPHIDE USING SAID CATALYST

of which the following is a full, clear and exact description.

A NOVEL CATALYST USEFUL FOR REMOVAL OF MYDROGEN SULPHIDE FROM GAS STREAM AND ITS CONVERSION TO SULPHUR, A PROCESS FOR PREPARING SUCH CATALYST AND A METHOD FOR REMOVING OF HYDROGEN SULPHIDE USING SAID CATALYST

#### FIELD OF THE INVENTION

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The invention relates to a catalyst comprising 0 to 95 % by weight clay, 0 to 95 % by weight gypsum and 0 to 95 % by weight alumina and 5 to 60 % by weight hydrated iron exide and heated to remperatures between 100 and 650°C for enhanced activity for removal of hydrogen sulphide from gas streams and its conversion to sulphur, a process for preparing such catalyst and a method for removing hydrogen sulphide using said catalyst.

# BACKGROUND AND PRIOR ART REFERENCES

Hydrogen sulphide is a highly toxic and corrosive environmental political with an obnexious smell which needs to be removed for polition control as well as process requirements in industries. Natural gas processing complexes, refineries, sulphur processing chemicals industries, pharmaceutical industries, sugar industries, sewage treatment plants and bio-gas generating units are some of the major industries which need an economically viable solution for H<sub>2</sub>S removal and its safe disposal.

A mainber of processes have been known and are in commercial use for removing hydrogen sulphide from gas streams. However, these processes have some inherent limitations. The processes used for removal of H<sub>2</sub>S and there disadvantages are described in detail hereafter.

Claus process is used for removing hydrogen sulphide from gases containing typically high concentration of H<sub>2</sub>S (more than 20% by vol of H<sub>2</sub>S). Liquid Redox process is used for removing hydrogen sulphide from gases containing typically low concentration of H<sub>2</sub>S. Both the aforesaid processes have the disadvantages of high capital and operating cost.

Processes using iron sponges as catalyst have been in use wherein from oxide deposited on wood shaving is used for removing hydrogen sulphide from gases. The major disadvantage with such a catalyst that these can be used as only once through catalyst i.e. the catalyst after being used for removal of H2S can not be regenerated and hence has to be disposed as waste. Therefore, the cost of such treatment is high this to the use of stoichiometric quantities of chemicals and also disposer of the used materials.

Further, loading capacity i.e. the extent upto which the wood shavings can be loaded with the iron oxide is low, due to which, the hydrogen sulphid removal capacity in a single pass is limited. Also, safe disposal of the used catalyst is major problem.

- In yet another process for hydrogen sulphide removal, a hot zinc exide hears used.

  Zinc exide is costlier than from exide. Another limitation of the process is that the bed gets exhausted after treating stoichiometric quantity of hydrogen sulphide ence through the bed. The need of higher temperature for effective removal is another disadvantage, as the gas needs to be preheated prior to treatment. Zinc exide gets converted to zinc sulphide which is disposed off after the bed gets exhausted.
- From the above descriptions of prior art, it is clear that there is a need for a more economical and simple process for hydrogen sulphide removal and its conversion to elemental sulphur using a solid bed incorporating inexpensive chemicals which can be regenerated and reused multiple times. This is the main objective of the present invention

# OBJECTIVE OF INVENTION

The objective of the present invention is to provide an iron oxide tused catalyst which can be used multiple times for removal of hydrogen sulphide from gas attenuit containing the same and its conversion to elemental sulphur

Another objective of the present invention is to provide process for preparing aforesaid

One another objective of the present invention is to provide a method for removal of sulphur compounds from a gas stream comprising the same and recovery of elemental sulphur therefrom using aforesaid catalyst.

## STATEMENT OF INVENTION

- The present invention relates to a catalyst useful for removal of hydrogen sulphide from gas streams containing the same and its conversion to elemental sulphin. He said catalyst comprising 0 to 95 % by weight clay, 0 to 95 % by weight gypsum and 0 to 95% by weight alumina and 5 to 60 % by weight hydrated from exide and heated to temperatures between 100 and 650°C (for enhanced activity and
- The present invention further relates to a process for preparing a catalyst useful for removing hydrogen sulphide from a gas stream and recovering elemental sulphur therefrom, said process comprising the steps of:

- a) mixing of 0 to 95 % by weight clay, 0 to 95 % by weight gypsum, 0 to 95 % by weight alumina and 5 to 60 % by weight hydrated iron oxide; and
- b) granulating, pelletizing or pulverizing the mixture of step (a) and heating the same at temperature in the range of 100°C to 650°C to obtain the catalyst.
- The present invention also relates to a method for removal of sulphur compounds from a gas stream comprising the same and recovery of elemental sulphur therefrom said method comprising the steps of:
  - a) mixing moist air/ water with the gas stream compounds for converting the sulphur compound to livdrugen sulphide.
  - a) Contacting the gas stream containing hydrogen sulphide with a catalyst comprising to 0 to 95 % by weight they; 0 to 95 % by weight gypoum; 0 to 95 % by weight alumina and 5 to 60 % by weight hydrated from oxide to remove hydrogen sulphide by chemisorption;
  - a) regenerating the spent chialyst by passing air through or over the same to oxides of iron and converting iron sulphides to from oxides and elemental sulphin.

#### SUMMARY

The solid material used for hydrogen sulphide is made by an inventive method to enable loading of the active content to high levels as well as improve its activity by a unique heat treatment method. The process also is designed to render the medium porous for greater gas penetration and availability of reactive sites. The repeated ability to regenerate the active chemical entity in the system renders the process catalytic in nature.

The chemical reactions which enable the process of hydrogen sulphide removal and regeneration of the active content of the solid medium are given below:

### A. Hydrogen Sulphide Removal Reactions

1. Fe<sub>2</sub>O<sub>3</sub> + 3 H<sub>2</sub>S Fe<sub>2</sub>S<sub>3</sub> + 3 H<sub>2</sub>O

2. Fe/S, 2 FeS | S

3 2 FeS + 1 1/2 O2 Fe2O3 + 2S:

4.  $3H_2S + 1 \% O_2 = 3S + 3 H_2O$ 

- B. Carbonyl Sulphide Removal Reactions
  - 5. 3COS + 3 H<sub>2</sub>Q: -- 3 CO<sub>2</sub> + 3 H<sub>2</sub>S
- C. Carbon Disulphide Removal Reactions
  - 6. CS2+2H2O -> CO2+2H2S
- Carbonyl sulphide and carbon disulphide are converted to hydrogen sulphide by reaction with water present with the treating gas or in the bed and the hydrogen sulphide produced is then converted into elemental sulphur as given above in equations.
- Iron oxide in the medium which is in the ferric oxide form reads with hydrogen sulphide to form ferric sulphide as shown in the first equation. Ferric sulphide being unstable gets converted to the more stable ferrous sulphide and sulphur (Eq. 2). During this process, iron gets reduced to ferrous form and hydrogen sulphide gets oxidized partially to sulphim. The ferrous sulphide on contacting with air yets oxidized as shown in the equation 3 to elemental sulphur and ferric oxide, thus regenerating the same for another cycle of reaction with hydrogen sulphide.
- This is thus a catalytic redox process wherein the ferric and ferrous forms of iron are formed during the reaction and regeneration cycles. The product of the reaction is elemental sulphur. The net result of the reaction cycle is the oxidation of hydringen sulphide to elemental sulphur by the oxygen in the air as shown in the equation 4
- Also, other sulphur containing compounds such as carbonyl sulphide and carbon disulphide are also convened into hydrogen sulphide as shown in reaction 5 & 6 and subsequently to elemental sulphur as given in Equations 1 to 4.

# DETAIL DESCRIPTION OF THE INVENTION

- The Present invention relates to a catalyst useful for removal of hydrogen sulphide trom gas streams containing the same and its conversion to elemental sulphur, the said catalyst comprising 0 to 95 % by weight clay, i) to 95 % hy weight gypsum and 0 to 95 % by weight hydrated iron exide and heated to temperatures between 100 and 650°C for enhanced activity:
- Yet another embodiment of the present invention, wherein the weight percentages of clay, gypsun and alumina are not simultaneously equal to zero.
  - Yet another embodiment of the present invention, wherein said catalyst comprising 5 to 60 % by weight gypsum and 5 to 40 % by weight alumina and 6 to 40 % by weight hydrated from oxide.

Yet another embodiment of the present invention, wherein chays are selected form the group comprising Kalonite, Montomorillonite/Someetite, Illite and Chlorite.

Yet another embodiment of the present invention, wherein clays are selected form the Semectite group.

Yet another embodiment of the present invention, wherein clay used is bentonite clay.

Yet another embodiment of the present invention, wherein said catalyst contains forrie

tons as active sites, which chemisorhs hydrogen sulphide present in the gas stream and

converts the same into elemental sulphur.

Yet another embodiment of the present invention, wherein said catalyst has pill value in the range of 8.0 to 10.0

Yet another embodiment of the present invention, whereus sant catalyst is meat treated at temperature in the range of 100° C to 650° C before use for activating the same.

Yet another embodiment of the present invention, wherein 100 gm of said catalyst chemisorbs 2860 to 28600 mg of hydrogen sulphide from said gas stream in one cycle.

5 Yet another embodiment of the present invention, wherein said spent catalyst containing sulphides of iron is regenerated by passing all through the same at ambient temperature.

Yet another embodiment of the present invention, wherein regenerated catalyst treats and tempores hydrogen sulphide, from the gas stream and converts the same to elemental sulphur in the subsequent cycles of chemisorption and regeneration.

Yer another embodiment of the present invention, wherein the catalyst is used in at least 15 chemisorption and regeneration cycles during its use.

Yet another embodiment of the present invention, wherein sulphides of iron present in the spent catalyst is converted to Fo<sub>2</sub>O<sub>3</sub> during regeneration thereby producing elemental sulphur and regenerating the catalyst.

You another embodiment of the present invention, wherein the elemental sulphur recovered has purity more than 99.%.

Yet another embodiment of the present invention, wherein said catalyst is used in fixed bed reactors or fluidized bed reactors.

Yet another embodiment of the present invention, wherein said entalyst is divided into fine particles having particle size in the range of 100 μm to 2000 μm for use in the fluidized had reactor.

Yet another embodiment of the present invention, wherein said catalyst is pelletized or granulat d to obtain pollets/ granules having diameter in the range of 0.5 nm to 10.0 mm for use in fixed bed seactors.

A further embodiment of the present invention relates to a process for preparing a catalyst useful for removing hydrogen sulphide from a gas stream and recovering elemental applicant therefrom said process comprising the steps of

- a) mixing of 0 to 95 % by weight clay 0 to 95 % by weight gypsum, 0 to 95 % by weight alumina and 5 to 60 % by weight hydrated iron oxide.
- b) granulating, pelletizing or pulverizing the mixture of step (a) and heating the same at temperature in the range of 100°C to 650°C to obtain the catalyst

Still further embodiment of the present invention, wherein it step (a), the hydrated from uxide is prepared from commonly available salts of irou such as forms nitrate, fearic chloride, feiric sulphate and commonly available alkali ammonium hydroxide, sodium hydroxide and potassium hydroxide.

Yet another embodiment of the present invention, wherein 100 gin of the catalyst thus obtained chemisorb 2860 to 28600 mg of hydrogen sulphide gas from the gas sceam.

Yet another embodiment of the present invention, wherein the catalyst thus obtained has pH value in the range of 8.0 to 10.0.

Yet another embodiment of the present invention, wherein the catalyst thus obtained is used in fixed bed reactor or fluidized bed reactor.

Yet another embodiment of the present invention, wherein catalyst thus obtained contain ferric ions as active sites.

Yet another embodiment of the present invention, wherein the catalyst thus ubtained is pulverized into fine particles for use in fluidized bed reactors.

Yet another embadiment of the present invention wherein in step (b), the mixture of step (a) is pelletized or granulated to obtain pellets/ granules having diameter in the range of 0.5 mm to 10 mm for use in fixed bed reactors.

Still further embodiment of the present invention relates to a method for removal of sulphur compounds from a gas stream comprising the same, and recovery of elemental sulphur therefrom, said method comprising the steps of:

a) mixing moist air/ water with the gas stream comprising the sulphur compounds for converting the sulphur compound to hydrogen sulphide:

- Contacting the gas stream containing hydrogen sulphide with a catalyst comprising to 0 to 95 % by weight clay, 0 to 95 % by weight gypsum, 0 to 95 % by weight alumina and 5 to 60 % by weight hydrated iron oxide to remove hydrogen sulphide by chemisorption, and
- c) regenerating the spent catalyst by passing air through or over the same to oxides of iron and converting from sulphides to iron oxides and elemental sulphur.

In yet another embodiment of the present invention, wherein compounds of sulphur are hydrogen sulphide, carbonyl sulphide (COS), and carbon disulphide (CS2) and mixtures thereof.

Yet another embodiment of the present invention, wherem the gas streams containing hydrogen sulphide from trace level to 100% level is treated to get outlet gas stream free of the same.

Yet another embodiment of the present invention, wherein the color of the establish changes from reddish brown to black during step (b) chemisorption and luchanges back to reddish brown on regeneration, this property being useful in visually monitoring the progress of the chemisorption and regeneration cycles respectively.

Yet another embodiment of the present invention, wherein the spent catalyst is regenerated by passing an oxygen containing gas through or giver the same.

Yet another embodiment of the present invention, wherein removal of the sulphur compound from the gas stream and regeneration of catalyst are optionally catried out simultaneously.

Still another embodiment of the present invention, wherein removal of the sulphur compound from the gas stream and regeneration of catalysters simultaneously carried out by contacting gas stream containing sulphur compounds & an oxygen containing gas simultaneously with the catalyst.

Yet another embodiment of the present invention, wherein the rate of simultaneous reaction and regeneration of catalyst depends on the flow rates of gas stream and ratio of gas stream and oxygen containing gas as well as the hydrogen sulphide content of the gas stream.

Vet another embodiment of the present invention, wherein the purcentage of regeneration of spent catalyst is 100% when oxygen containing gas is passed through on over the spent catalyst.

Yet another embodiment of the present invention, wherein the process is carried out in fluidized bed reactors or fixed bed reactors.

Yet another embodiment f the present invention, wherein the elemental sulphur obtained has purity more than 99 %.

Yet another embodiment of the present invention, wherein 100 gm of said catalyst chemisorbs 2860 to 28600 mg of hydrogon sulphide from said gas stream in one cycle. The invention is different from the ones reported so far as that a solid medium incorporating iron hydroxide in the bulk of the same is prepared by mixing the ingredients which are naturally occurring, non toxic and non hazardous in natura with iron hydroxide and heat treating the same to get high activity for hydrogen sulphide removal and its conversion to elemental sulphur. The iron hydroxide is prepared from any common iron salts such as iron chloride, iron sulphate and iron nurate by treatment with alkalis such as sodium hydroxide; potassium hydroxide or ammonium hydroxide. The mixture of iron hydroxide and the support medium is converted to granules or 15 pellets for easy packing (filling) in a column and treated to temperatures between 1111 to 600°C to increase the reactivity of the iron oxide towards hydrogen sulpitide as well as to make the regeneration of the same with oxygen containing gases possible. The granules help to reduce pressure drop across the column through allowing easy passing of the gas through the same. This eliminates requirement of high pressure for the gas being treated.

The iron hydroxide in the medium is converted to iron axide by a process of heat treatment of the granules / pellets: Preparation of the solid medium incorporating the iron sail is done at relatively low temperatures as compared to the one reported recently (ref. US Patent 6500237) wherein a calcined material is used for impregnation of the active matter wherein the iron hydroxide adheres to the exposed surfaces of the medium. The total hydrogen sulphide treatability is also found to be higher as compared to the prior art.

Another advantage of the process is that the sulphur deposited on the solid medium can be recovered by extraction with a suitable solvent like carbondsculphide or carbon tetrachloride or other organic solvents in which sulphur is soluble. Sulphur can also be extracted by heating the medium above sulphur melting temperature as a solid or alternately by sharing in water and heating the slurry to above the melting point of sulphur. The molten sulphur can be separated from the slurry containing the support

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medium. The recovered sulphur is of high quality and can be used for dewnstream applications. In cases where the user is not interested in extraction of sulphur the bed can be disposed off safely without further treatment due to the non-toxic nature of the medium and its contents.

We thus report here an improved process for hydrogen sulphide removal from this streams using a novel solid bed made by incorporating from oxide in a mixture of materials and heating the same to a temperature high enough to make it chemically active and porous for easy availability of the reactive sites in the solid. The material can be regenerated using a simple process and reused multiple number of times in convent hydrogen sulphide to elemental sulphin. The sulphur thus deposited on the bed can be recovered using methods known in prior art.

Accordingly, the present invention provides a novel catalytic process for hydrogen, sulphide removal from sour gas streams and its conversion to sulphur using regenerative solid bed consisting of finely divided from oxide or its hydrated form made from common salts of iron such as chloride, sulphate and nitrate and an alkali such as bydroxides of sodium, potassium or ammonium and incorporated in a support medium consisting of naturally occurring clays and minerals singly or as mixture to impart stability to the granules or pellets made from the mixture followed by heating the granules or pellets to a temperature high enough to enhance its reactivity towards hydrogen sulphide as well as enabling the regeneration of the iron oxide by conversion of the iron sulphide formed to sulphur and iron oxide on treatment with oxygon containing gas, the size of the pellets or granules being not limiting in the hydrogen sulphide removal characteristics of the solid bed medium.

The Applicant surprisingly found that in the composition of the catalyst, the amount of iron exide incorporated plays a vital role in determining the suitability of the catalyst in the process of removal of hydrogen sulfide from the gaseous stream. More particularly, the applicants noticed that if the amount of iron exide is incorporated in the catalyst composition was less than 5 % by weight, the catalyst did not efficiently remove H<sub>2</sub>S from the gaseous stream. The applicants were of the opinion that increasing the amount of hydrated iron exide in the catalyst composition would increase its efficacy in removing H<sub>2</sub>S from the gaseous stream. However, surprisingly the applicants above hypothesis were found to be wrong. The applicants surprisingly naticed that increasing the amount of hydrated iron exide incorporated in the catalyst composition beyond a

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certain range adversely affected other properties of the catalyst and made it insultable for us in the process. Mor particularly, increasing the amount of the hydrated iron exide incorporated in the natalyst composition beyond 60 % adversely affected the pelletization and granulizing properties of the catalyst. As the main aim of the present invention is to provide catalysts which are stable enough for regeneration, any adverse effect on the pelletization and granulization properties of the catalyst rendered the same unsuitable for even a single regeneration.

The applicants would also like to cumbiasis here that in step of heating the catalyst prior to use plays a vital role on the efficacy of the process for removal of H2S from the gaseous state. The applicant noticed that if the catalyst is used without prior heating, the removal of H2S content from a gaseous stream is not significant. This is due to the fact that pore formation in the catalyst does not take place and hence very less contact surface area is svailable for the observation of H2S gas. Applicant also noticed that if the catalyst is ficated prior to use, pores are developed in the catalyst and enhance the absorption of the H2S gas by providing more contact surface area.

Applicant also noticed that if the catalyst is heated prior to use, some from exide present in the interior part of the catalyst, come out on the outer surface and provide enhanced activity to the catalyst.

Applicants also noticed that leading the catalyst continuously and above a certain temperature adversely affects the activity of the catalyst. More particularly, the applicants notice that the heating the catalyst above 600°C desmoy the catalytic activity, the applicants found that when the catalyst is heated above the 600°C the man oxide undergoes a transformation in the state and and the transformed state does not provide any catalytic activity.

Hence, the amount of iron exide included in the catalyst and the temperature up to which the catalyst is heated are critical and non-obvious aspects of the interesting invention. None of the document available, teach or suggest the these critical and non-obvious features.

This invention is described in detail in the following examples which are provided by way of illustration only and therefore should not be construed to limit the scope of the invention.

### BRIEF DESCRIPTION OF TABLES

Table I shows the results obtained of hydrogen sulphide removal from a gas stream at various gas flow rate.

Table 2 compares the result obtained of hydrogen sulphide removal from a gas stream for a heat treated catalyst with non-heat treated catalyst.

Table 3 shows the result obtained of hydrogen sulphide removal from a gas stream mix with N g or CO2 or CH2 and sir.

Table 4 shows exygen content in outlet gas stream after passing through said catalyst.

Table 5 shows number of regeneration cycle performing for hydrogen sulfide removal with said catalyst.

Table 6 shows result obtained of hydrogen sulphide removal from a gas stream having various H2S: O2 ratio.

#### **EXAMPLES**

# Example 1

- 15 A solution of iron (III) nitrate (1000 g) in water is prepared and was freated with sodium hydroxide solution (20 g in 100 g water) in an agitated vessel to precipitate itum hydroxide. The precipitated iron hydroxide was allowed to settle, the supernatura clear liquid was discarded and the solid recovered by filtration and washed with water to remove disserved salts:
- The iron hydroxide (250 gm) thus isolated was mixed thoroughly with the solid support material bentonite clay (250 gm), alumina (125 gm) and gypsum (700 gm) and converted to granules (3mm diameter) in a granulating drum or pellets in a pelletiser (4 mm diameter).
  - The granules/pellets were dried, treated at temperature of 450 to 550 °C and used for removal of hydrogen sulphide and other toxic gases contained in gas streams as given in the following examples:

# Example 2

A solution of iron (III) nitrate (1000 g) in water is prepared and was treated with sodium hydroxide solution (20 g in 100 g water) in an agitated vessel to precipitate iron hydroxide. The precipitated iron hydroxide was allowed to settle, the supernatural clear liquid was discarded and the solid recovered by filtration and washed with water to remove dissolved salts.

The wet hydroided from hydroxide obtained above (560 gm; corresponding to 11.50 % ferric hydroxide on dry basis) was mixed th roughly with the solid support material bentonite clay (250 gm), alumina (125 gm) and gypsum (700 gm) and converted to granules (3mm diameter) in a granulating drum or patiets in a pelicuser (4 mm diameter).

The granules/pellets were dried, treated at temperature of \$50 to \$50 °C and used for removal of hydrogen sulphide and other toxic gases contained in gas streams as given in the following examples. The pellets thus obtained had iron content of 6.0 % by will and good granule integrity and crushing strength:

# 10 Example 3

The wet hydrated iron hydroxide obtained above (1500 gm, corresponding to 68 % ferric hydroxide on dry basis in the mixture) was mixed characally with the solid support material bentonite clay (100 gm), alumina (50 gm) and gypsum (125 gm). The material was granulated in a granulator, however, it could not be formed but granules of good trushing strength. Attempts at palletisation also failed

### Example 4

The solid bed medium (225 gms); reddick brown in colour prepared as given in Example I above was packed in a glass column of 32 mm diameter and 350 mm height. Gas containing a mixture of hydrogen sulphide (1.14 % by volume), and rest phrogen was passed through the bed at the flow rate of 0.30 line perindicute. The pattlet gas was found to be free from hydrogen sulphide. The bed became black in colour as the hydrogen sulphide reacted with it and when the bed was exhausted, the material became totally black as shown by the prescue of hydrogen sulphide in the outlet gas. Heat generation was observed during the chemisorption cycle.

Through the medium in the column which was now black in colour, ambient sir was passed. Slowly the column restored to its original reddish brown colour, thus indicative of its regeneration. Heat generation was observed during the regeneration cycle.

Through the above regenerated medium, hydrogen sulphide containing gas was again passed as above and the outlet gas was found to be free from the sour gas. The reaction regeneration cycle was repeated 20 times in this manner and the cultumit was found to be reactive to hydrogen sulphide removal without significant reduction in hydrogen sulphide removal without significant reduction in hydrogen sulphide removal capacity.

Example 5

As described in example 2 above, gas containing a mixture of hydrogen sulphide (4.7 %, 47000 ppm by volume) and rest nitrogen was passed through the bed at the flow rate of ():14th litre per minute. The outlef gas was found to be free from hydrogen sulphide. The bed became black in colour as the hydrogen sulphide reacted with it and when the bed was exhausted, the material became totally black as evident by the presence of hydrogen sulphide in the outlet gas.

#### Example o

The solid bed medium 100 g as prepared in example 1 was taken in a glass nothing of 32 mm diameter. Catalyst bad height measured which was 14 centimeters. Maist Clascontaining a mixture of hydrogen sulphide (15.4 % by volume) and rest nitrogen was passed through the bed at the flow rate of 0.230 litre per minute. The mixing was passed till hydrogen sulphide concentration in the outlet gas stream reached 100 ppm.

# Example 7

The solid hed medium (150 gms), reddish brown in colour, prepared as given in Example I above, was packed in a glass column of 32 mm diameter and 235 mm height. Pure hydrogen sulphide was passed through the bed at the flow rate of 0.04 line per minute. The bed became black in colour as the hydrogen sulphide reacted with it. Outlet of the column was passed through a cadmium acetate solution (1 gm cadmium acctate dissolved in 100 gms of water) to detect hydrogen sulphide presence in the treated gas. (Hydrogen sulphide reacts with cadminum acetate to form yellow precipitate of cadmium sulphide). Hydrogen sulphide was not detected in column outlet until last 2 cms of unexhausted bed was available for reaction with hydrogen sulphide as per the visual indication. Heat generation was observed during the chemiscription cycles Example 8

# The solid bed medium, 100 g, as prepared in Example I was tilled in a glass column of 32 mm diameter to a height 14.5 centimeters. Gas containing a mixture of hydrogen sulphide (3 % by volume) and rest methane was passed through the bed at the flow rate of 0.20 litre per minute. The results of outlet stream hydrogen sulphide concentration noted and are as given below.

After passing 37 litres of gas in 185 min, 100 ppm of hydrogen sulphild was observed. 30 in the outlet gas and the column was taken for regeneration. After passing ambient air through the bed, it regained its original colour and became active for next cycle of chemisorption.

# Exampl 9

The solid bed medium (100 g), reddish brown in col ur prepared as given in Example I shove was packed in a glass column of 32 mm diameter. Gas commining a mixture of hydrogen sulphide (9.1 % by volume), and rest carbon dioxide was passed through the bed at the flow rate of 0.05 line per minute. As the hydrogen sulphide reacts with the ferric ions to form iron sulphide, the colour of the bed changes from reddish to black. Hydrogen sulphide in treated stream was found to be helow tracaphle level until 15.86 litres of gas mixture was passed. The experiment was continued fill hydrogen sulphide concentration in treated gas stream and inlet gas stream became same. Total 40.26 hires of gas was treated in this manner.

Through the solid bed in the column which was now black in colour, ambient moist air was passed till it regained its original colour, indicative of its regeneration. Hour generation was observed during the regeneration cycle.

# Example 10

The solid had medium (100 g), reddish brown in colour prepared as given in Example I above is packed in a glass column of 32 mm diameter. Gas containing a mixture of hydrogen sulphide (4.75 % by volume) with nitrogen was passed through the column. The experiment was repeated under identical conditions with frash had, but with different gas flow rates. In all cases gas was passed till hydrogen sulphide level in outlet gas stream reached 100 ppm level. Quantity of gas treated and hydrogen sulphide removed were measured Results are as given in Table 1 below.

Table I

Gas Flow	Gas Velocity	Total volume.	Volume of	Residence
Rate		of gas treated :	hydrogen sulphide	Time;
	M/min	(Litres)	removed. Litres	Seconds
ml/min				
1.00	0.12	19:53	0.93	62.70
150	0.18	22.275	1.06	41.80
300	0.36	18.36	0.87	21 70
400	0.48	7.20	0.34	16.64

As the volume of gas that could be treated remained nearly same upto a flow rate of 300 ml/ min, the above example has shown that the residence time required for reaction was about 21 secs.

#### Example 11.

The solid bed medium (100 g), reddish brown in colour prepared as given in Example 1 shove was packed in a glass column of 32 mm diameter. Gas containing a mixture of Hydrogen sulphide (5 % by volume) with nitrogen was passed through the column: Keeping the gas mixture same, experiment was conducted for following continuous.

- Solid bed medium treated with ambient moist air for I hour before treatment with gas. Moist gas was passed through the catalyst bed till hydrogen sulphide concentration in the outlet gas stream reached above 100 ppm.
- ii. Solid bed medium treated with ambient moist air for I ligur before treatment with gas. Dry gas was passed through the solid bed till hydrogen aulphide concentration in the outlet gas stream was more than 100 ppm.
- iti. The catalyst was not given any treatment prior to experiment. Dry gas mixture was passed through the solid bod till hydrogen sulphide concentration in the outlet gas stream was more than 100 ppm.
- 15 The results of above three cases are given in Table 2 below.

Table 2

Hydrogen Sulphide Concentration : 5 % by volume

	Gas Velocity	Residence	Total gas treated	Total H2S
	Cm/Second	Time Seconds	until outlet H-S	removed
			concentration	
			reached 100 ppm	Litres
			Litres	
Case I	0.41	31.36	18.00	0.90
Cose II	0.41	31.36	18.90	0.945
Case III	0.41	31.36	14.40	0.72

The results show that in case where the bed was pre-treated with moist ambient hir, higher hydrogen sulphide removal capacity was observed compared to the hed which was not given pre-treatment with moist ambient air. Moisture content in the bed and for moisture in the gas were found to improve hydrogen sulphide removal efficiency.

# Example 12

The solid bed medium (225 g), reddish brown in colour prepared as given in Example I above was packed in a glass column of 30 mm diameter, bed height of 34 contimeters. Gas containing a mixture of carbon disulphide, (35 ppm by volume) with carbon

dioxide was passed through the column at a flow rate of 50 in per minute. Carbon disulphide (CS<sub>2</sub>) concentration in the treated gas was measured and found to be below traceable levels.

Example 13

- The solid bed medium, reddish brown in colour prepared as given in Example I above was packed in a glass column of 15 mm diameter, catalyst bed height of 25cm. Casseontaining a mixture of carbonyl sulphide, COS (5 ppm by volume) with hitrogen was passed through the column at ambient temperature. Carbonyl sulphide (COS) was not tracepble in the treated gas.
- (i) Example 14

The solid catalyst (25g) which underwent 8 chemisorption and regeneration cycles was taken in a closed vessel and was mixed with 75g of water. The mixture was healed at 1.25° C for 30 minutes. It was found that sulphur contained in the catalyst melted and separated out from the rest of the solid under these conditions. The vessel was cooled down and hunps of sulphur were recovered.

Example 15

Through the solid catalyst bed (25 g) which had undergone 8 chemisorpion and regeneration cycles packed in a glass column, Carbon disulphide (CS<sub>2</sub>) was passed from the top at the flow rate of 20 ml per minute. As CS<sub>2</sub> passed through the column, it dissolved the sulphur and the sulphur containing solution was collected at the bottom. Sulphur extraction was continued for 1 hour in this manner. From the CS<sub>2</sub> solution containing sulphur, CS<sub>2</sub> was distilled out and sulphur, bright yellow crystalline, was isolated. Sulphur thus obtained had a purity of 99.99%.

Example 16

- The solid bed medium (100 g), reddish brown in colour prepared as given in Example 1.

  above is packed in a glass column of 32 mm diameter and 130 mm height. The
  following gas mixtures were passed through fresh polumns of identical dimensions with
  gas flow rate of 0.2 litres per minute in all cases, monitoring the outlet gas quality.
  - llydrogen sulphide (4.4 % by volume) with Carbon dloxide.
- 10 ii. Hydrogen sulphide (5.1 % by volume) with nitrogen,
  - iii. Hydrogen sulphide ( 4.26 % by volume) with methane and
  - iv. Hydrogen sulphide (4.25 % by volume) with air

It was found that in all cases, the hydrogen sulphide concentration in outlet was helow detectable limits. One was passed until hydrogen sulphide concentration in number gas stream reached 100 ppm by volum. The quantity of gas trated in the first pass is given in the table below with other parameters.

5 Quantity of gas treated and hydrogen sulphide removed is given in Table 3 below.

Table 3

Sr.No.	Gas Mixture	Duration 7	Gas Flow	Total Gas	Total H2S
•		Minutes	l per nun	treated, I	Removed,
					Litres
1 : .	Nitrogen +	100	0.20	18	0.92
	H <sub>2</sub> S(5.11%)				
2	Methane +	125	0.20	22.5	0.96
	H <sub>2</sub> S(4.26%)				
3	Carbon	115	0.20	20.7	.0.91
	Dioxide +				
	H <sub>2</sub> S(4,40%)				
4	Air.+	835	0,30	225.5	9.58
	H <sub>2</sub> S(4.25%)			Lord And	E SE PO

Above data indicates that hydrogen sulphide removal capacity is not significantly affected by the presence of carbondioxide, nitrogen and methane. In case of air chemisorption and regeneration were found to take place simultaneously, and the best could be used continuously, thus resulting in the treatability of about 10 times the gas that could be treated in one pass in the absence of air.

#### Example 17

15

The solid bed medium (229 g) which was heated with hydrogen sulpline (black in colour) was packed in a glass column of 30 mm dia. Height of the catalyst bed was 300 mm. Catalyst which was black in color was treated with ambient ait for regeneration. Air at the rate of 0.1 Litres por minute was passed through the column. Catalyst colon started changing from black to grayish and ultimately to reddish brown with yellow tinge. Color change started from the hoftom portion and moved up as bottom layer of catalyst got regenerated. Outlet air samples were analyzed for oxygen content. The results are given in Table 4 below.

Table 4

Sr.No. Time (			( Minutes)	Oxygen Content %
1	٠	<del>.</del>	0	20.8
2	· ; ;	: ' : .	30	4 84
· 3	:	• • • •	60	5:32
4		77.11	90	8.46
5	:		150	17,35
. 6			200	20.5

Regeneration was found to be complete after 200 minute and the hed revained hydrogen sulphide removal activity. During regeneration, heat generation was observed and moisture was found deposited on the walls of the column.

### Example: IX

The solid bed medium (225 g), reddish brown in colour prepared as given in Example 1 above is packed in a glass column of 32 mm diameter. Catalyst bed height was 310 mm. Gas containing hydrogen sulphide and carbon dioxide was passed from the bottom of the column. After exhaustion of the catalyst heil, the same was regenerated with ambient air. Regenerated catalyst bed was again used for hydrogen sulphide removal. Reaction and regeneration cycles were carried out on the same column 20 times even after which the hert maintained its hydrogen sulphide removal capacity. The colour of the bed became yellowish due to presence of elemental sulphur. Results of the different runs are given in Table 5 below.

Tal

	٠.					:			<u> </u>	المناخف
Run No.	1	2	.3	4	5	.6	7	8	9	10
H <sub>2</sub> S Treated, lit	4.01	3.91	3.59	3.64	2.75	3.38	4.35	4.87	3.55	3.49
Run No	11	12	13	14	.15	16	17	18	19	20
H <sub>2</sub> S.Treated, lit	3.92	4.16	3.73	1:. 1						

Example 19

The solid bed medium (100 g), reddish brown in colour, prepared as given in Example 1 above, was taken in the glass column of 32 mm dia and the following experiments were conducted with gas and air mixtures.

Case I. A gas mixture containing hydrogen sulphide (1 % by volume in carbondioxide) and air were mixed in the ratio of 1 0.075 ( .075 liter air per litre of gas rand the mixture at a flow rate of 0.30 litres per minute was passed through the column, to

enable concurrent reaction and regeneration cycles. As oxygen is the regeneration agent, the gas mixture was prepared such that hydrogen sulphide to oxygen ratio was around 1: 1.5. The treated gas was tested for the presence if hydrogen sulphide: Hydrogen sulphide in our let gas stream was not traceable. The results and observations are given in the Table 6 below.

Case H: A gas mixture containing hydrogen sulphide (1.5 % by volume in carbondioxide) and air were mixed in the ratio of 1: 0.30 (0.30 litre air per litre of gas) and the mixture at a flow rate of 0.30 litres per minute was passed through the column, to enable consurent reaction and regeneration bycles. As oxygen is the regeneration agent, the gas mixture was prepared such that hydrogen sulphide to oxygen ratio was around 1: 4. The treated gas was tested for the presence of hydrogen sulphide. Hydrogen sulphide in out let gas stream, was not traceable. The results and observations are given in the Table 6 below.

Table 6

	H <sub>2</sub> S Conc.%	H2S: O2 Ratio	Gas Flow Rate	fotal Clas		
			Li/Minutes	Treated 1		
Case I	1:	I:1.5	0.300	64.23		
Case II	1.5	1 4	0:300			

Results of the above experiments indicate that with H2S to axylien ratio higher than 1:

4, the hydrogen sulphide removal and its conversion to sulphin can be carried out in a single step without separate reaction and regeneration cycles. This is particularly useful for cases where the gas containing hydrogen sulphide has no downstream applications, hence can be treated for its removal and then vented to air.